Boehringer Ingelheim is pleased to provide you with this Educational Pack, which has been developed to give practical and relevant information on the appropriate use of Pradaxa®.

The pack includes:

- Pradaxa® 110mg Summary of Product Characteristics
- Pradaxa® 75mg Summary of Product Characteristics
- Prescriber Guide this addresses recommendations for the use of Pradaxa® in order to minimise the risk of bleeding
- Patient Alert Card

To order additional copies of the Patient Alert Card please go to: www.pradaxa.co.uk/pVTEeducationalpack

You can also order or download this Educational Pack.

Reference

Boehringer Ingelheim. Pradaxa** 110mg hard capsules Summary of Product Characteristics.
 Boehringer Ingelheim. Pradaxa** 75mg hard capsules Summary of Product Characteristics.

Prescribing Information (pVTEp UK) – PRADAXA® (dabigatran etexilate)

Capsules containing 75 mg or 110 mg dabigatran etexilate (as mesilate) **Action:** Direct thrombin inhibitor indication: Trimary prevention of venous thromboembolic events is adult patients who have undergone elective total hip or knee replacement surgery. **Dos and Administration:** Renal function should be assessed by calculating CrCL prior to initiation to exclude patients with severe renal impariment (CrCL < 30 mL/min). Recommended dose is 220 mg once daily orally taken as 2 capsules of 110 mg. Initiat treatment within 1-4 hours of completed surgery with a single capsule continuing with 1 capsules once daily for a total of 10 days, (lone replacement surgery) or 28 – 35 day (hip replacement surgery). Delay initiation of treatment if haemostasis is not secured. I treatment is not started on the day of surgery, initiate with 2 capsules once daily, For the following groups the recommended daily dose of Pradaxa is 150 mg taken once daily a 2 capsules of 75 mg: patients with moderate renal impariment (CrCL 30-50 mL/min) patients who receive concomitant verapamil, amiodarone, quinidine; patients aged 75 o above. In patients with moderate renal impariment and concomitant verapamil, conside 75mg daily, Pradaxa is contraindicated in severe renal impariment (CrCL < 30 mL/min) Assess renal function by calculating CrCL prior to initiation to exclude patients with severe rena impariment. Renal function should also be assessed while on treatment in certai clinical situations when it is suspected that renal function could decline or deteriorate Not recommended if liver enzymes > 2 Upper Limit of Normal (ULN). No dose adjustmen required but close clinical surveillance in patients <50 kg or >110 kg. If switching fror Pradaxa to parenteral anticoagulant to Pradaxa, discontinue the parentera niticoagulant and start Pradaxa 0-2 hours prior to the time that the next dose of thatemate therapy would be due, or at the time of discontinuation in case of continuou treatment. No relevant uses of Pradaxa; a the patients should be instructe

platelet defects, recent biopsy, major trauma, bacterial endocarditis, oesophagitis, gastritis or gastroesophageal reflux. Concomitant use of ticagreior. The measurement of dabigatran related anticoagulation may be helpful to avoid excessive high exposure to dabigatran in the presence of additional risk factors. Patients who develop accute renal failure must discontinue Pradaxa. If severe bleeding occurs, discontinue treatment and investigate the source of the bleeding. Avoid or use with caution medicinal products which may increase the risk of haemorrhage. Avoid concomitant administration with P-gp inducers. Patients on dabigatran etexilate who undergo surgery or invasive procedures are at increased risk for bleeding therefore surgical interventions may require the temporary discontinuation of dabigatran etexilate, prescribers should consult the Summary of Product Characteristics for further information. Procedures such as spinal anaesthesia may require complete haemostatic function. The risk of spinal or epidural haematoma may be increased in cases of traumatic or repeated puncture and by the prolonged use of epidural catheters. After removal of a catheter, an interval of at least 2 hours should elapse before the administration of the first dose of dabigatran etexilate; these patients require frequent observation for neurological signs and symptoms of spinal or epidural haematoma. Treat with caution patients at high surgical mortality risk and with intrinsic risk factors for thromboembolic events. No data on the use of Pradaxa in patients undergoing hip fracture surgery, therefore treatment not recommended. Contains Sunset Yellow (E110) which may cause allergic reactions. Interactions: Anticoagulants and antiplatelet aggregation medicinal products; P-gp inhibitors coadministration (close clinical surveillance); amiodarone, quindine, verapamil reduce Pradaxa doss to 150mg (see above); consider dose reduction to 75 mg daily in patients with both moderate renal impairment and on verapamil; close monitoring with

Adverse events should be reported. Reporting forms and information cabe found at www.mhra.gov.uk/yollowcard. Adverse events should also be reported to Beetringer Incolleging Prus Safety on 0900 229 1627 (freeshort).



Date of preparation: June 2014 Job code: UK/DBG-141126

PRADAXA® (DABIGATRAN ETEXILATE) EDUCATIONAL PACK

For primary prevention of venous thromboembolic events in adult patients who have undergone elective total hip replacement surgery or total knee replacement surgery^{1,2}